APPLICATION FOR LETTERS PATENT OF THE UNITED STATES OF AMERICA

For the invention entitled:

CALCIUM HYPOCHLORITE OF REDUCED REACTIVITY

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CALCIUM HYPOCHLORITE OF REDUCED REACTIVITY

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3 FIELD OF THE INVENTION

- 4 This invention relates to calcium hypochlorite blends;
- 5 particularly to blends of solid calcium hypochlorite in
- 6 combination with an alkali salt form(s) of anti-scaling
- 7 polymers, wherein the polymer(s) are present in an alkali
- 8 salt form effective at reducing the hygroscopic
- 9 characteristics of the solid polymer(s), and subsequent
- 10 calcium hypochlorite blend. This invention also relates to a
- 11 means of reducing the reactivity of solid calcium
- 12 hypochlorite with the said polymer(s) by applying a coating
- of the polymer(s) to the solid calcium hypochlorite.

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BACKGROUND OF THE INVENTION

- 16 U.S. Patents 5,112,521 and 5,004,549 disclose the
- 17 blending of Phosphono-Butane-Tricarboxylic acid (PBTC) with
- 18 calcium hypochlorite to produce a solid calcium hypochlorite
- 19 blend that inhibits scale formation associated with the
- 20 reconstitution and dispensing of the calcium hypochlorite
- 21 solution.
- U.S. Patent 6,146,538, the contents of which are herein
- 23 incorporated by reference, discloses the use of Polymaleic
- 24 acid (PMA) and/or Polyepoxysuccinic acid (PESA) in an alkali

- 1 salt form that is blended with calcium hypochlorite to
- 2 reduce or eliminate scale formation associated with the
- 3 reconstitution and dispensing of the calcium hypochlorite
- 4 solution.
- It is known that increased moisture, especially
- 6 localized moisture, increases the rate of decomposition of
- 7 calcium hypochlorite. During decomposition, oxygen is
- 8 released, with chlorides and chlorates typically being the
- 9 resulting byproducts. In the presence of sufficient
- 10 moisture, exothermic reactions result, oxygen release
- 11 increases, and a potential fire hazard is produced when fuel
- 12 sources are available, e.g. cloth, hydrocarbons and the
- 13 like.
- 14 PBTC disclosed in U.S. Patents 5,112,521 and 5,004,549,
- 15 when combined with calcium hypochlorite as an anti-scaling
- 16 agent, can nevertheless compromise manufacturing safety and
- 17 solid product stability, due to its inherent hygroscopic
- 18 characteristics.
- 19 It has now been discovered that the alkali salt forms
- of PMA, PESA, and similar polymeric salts which are commonly
- 21 employed as deposit control agents, can significantly
- 22 reducing the hygroscopic characteristics of calcium
- 23 hypochlorite when combined therewith, in addition to
- 24 providing enhanced anti-scaling properties. It has been

- 1 further discovered that a coating of these polymers or
- 2 blends thereof significantly reduces the reactivity of the
- 3 solid calcium hypochlorite blend.
- 4 Thus, a step forward in the art is realized via
- 5 provision of a solid calcium hypochlorite combination which
- 6 simultaneously exhibits anti-scaling properties and reduced
- 7 hygroscopic and reactivity characteristics. Such a
- 8 composition yields improvements in both manufacturing safety
- 9 and product stability by reducing moisture adsorption and
- 10 subsequent decomposition of the calcium hypochlorite, as
- 11 well as improved safety by reducing the reactivity of the
- 12 solid calcium hypochlorite.

14 DESCRIPTION OF THE PRIOR ART

- U.S. Pat. No. 4,146,676 describes calcium hypochlorite
- 16 particles coated with about 4 to 45 percent by weight of a
- 17 low melting inorganic salt, e.g. aluminum sulfate hydrates,
- 18 to reduce dusting during handling and to increase stability
- 19 of calcium hypochlorite contacted with lighted cigarettes or
- 20 reactive organic materials such as glycerine.
- U.S. Pat. No. 4,965,016 describes a granular calcium
- 22 hypochlorite composition comprising a particle of calcium
- 23 hypochlorite, the surface of which has been covered with a
- 24 double decomposition product of calcium hypochlorite and an

- 1 alkali metal hydroxide. It was shown the granular strength
- 2 and stability was enhanced.
- 3 None of the prior art describes addition of an alkali
- 4 salt of an organic polymeric anti-scaling agent to provide
- 5 improved deposit control, while reducing the stability and
- 6 reactivity of the solid calcium hypochlorite.

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SUMMARY OF THE INVENTION

- 9 The instant invention is directed toward a solid
- 10 calcium hypochlorite product formulation and a method for
- 11 its production, which product exhibits improved
- 12 environmental stability and shelf-life while simultaneously
- 13 inhibiting scale in dispensing equipment, as well as in and
- 14 on feed equipment used for delivering a calcium hypochlorite
- 15 solution to a water stream which is to be treated.
- A solid calcium hypochlorite composition with improved
- 17 hygroscopic and reactivity characteristics is formulated
- 18 comprising hydrated calcium hypochlorite and an alkali salt
- 19 of Polymaleic acid, Polyepoxysuccinic acid or mixtures
- thereof, having between 0.01 and 10% by weight of said
- 21 alkali salts, an available chlorine concentration of at
- least 30% and water in the range of about 2% to about 20% by
- 23 weight.
- In a particularly preferred embodiment, the composition

- 1 is a solid having greater than about 50% active Calcium
- 2 Hypochlorite which exhibits anti-scaling characteristics and
- 3 reduced hygroscopicity and reactivity.
- 4 The solid calcium hypochlorite may be coated and/or
- 5 encapsulated with at least one polymeric alkali salt
- 6 selected from within the group including polymaleate,
- 7 polyacrylate, polyacrylamide, polycarboxylate,
- 8 polymethacrylate, Phosphinopolycarboxylate, Carboxylate-
- 9 Sulfonate copolymer, Maleic Anhydride copolymer,
- 10 Polyepoxysuccinate, maleate-sulfonate copolymer, maleate-
- 11 phosphonate copolymer, carboxylate-phosphonate copolymer,
- 12 and/or mixtures thereof.
- The alkali salt may be selected from at least one of
- 14 the group including the sodium, potassium, lithium, calcium,
- 15 or magnesium salts and mixtures thereof. Additionally, the
- 16 polymeric alkali salt may be in a liquid, slurry, or solid
- 17 form.
- 18 At some point in the manufacturing process the calcium
- 19 hypochlorite may be formulated/treated with a deposit
- 20 control agent including at least one alkali salt(s), e.g.
- 21 sodium, potassium, lithium, calcium, magnesium and/or
- 22 mixtures thereof, of polymaleic acid, polyexpoxysuccinic
- 23 acid, maleic anhydride copolymer, phosphinopolycarboxylic
- 24 acid, carboxylic-sulfonic acid copolymer, maleic-sulfonic

- 1 acid copolymer, maleic-phosphonic acid copolymer,
- 2 carboxylic-phosphonic acid copolymer, and/or mixtures
- 3 thereof. The weight percent of deposit control agent is
- 4 within the range of about 0.01% to about 10% of the final
- 5 weight of the formulated calcium hypochlorite product.
- 6 These polymers and copolymers will exhibit similar
- 7 hygroscopic characteristics in the desired salt forms while
- 8 providing deposit control capability. Furthermore, all are
- 9 hydrophilic and possess high thermal stability. Therefore
- 10 they will reduce reactivity of the solid calcium
- 11 hypochlorite product.
- 12 Accordingly, it is an objective of the instant
- 13 invention to provide a solid calcium hypochlorite product
- 14 and a method for its production, wherein the resultant
- 15 product exhibits reduced hygroscopic characteristics thereby
- 16 improving the manufacturing safety and shelf-life and anti-
- 17 scaling properties of the solid product.
- It is a further objective of the instant invention to
- 19 provide a coated or encapsulated solid calcium hypochlorite
- 20 product having reduced reactivity and anti-scaling
- 21 properties.
- Other objects and advantages of this invention will
- 23 become apparent from the following description taken in
- 24 conjunction with the accompanying drawings wherein are set

- 1 forth, by way of illustration and example, certain
- 2 embodiments of this invention. The drawings constitute a
- 3 part of this specification and include exemplary embodiments
- 4 of the present invention and illustrate various objects and
- 5 features thereof.

7 <u>DETAILED DESCRIPTION OF THE INVENTION</u>

- 8 As illustrated in Table 1, embodiments disclosed in
- 9 U.S. Patents 5,112,521 and 5,004,549 as well as an alkali
- 10 salt form of the Polymaleic acid disclosed in U.S. Patent
- 11 6,146,538 (all of which are herein incorporated by
- 12 reference) were tested for their hygroscopic properties by
- 13 exposing similar quantities of each to a controlled
- 14 atmospheric environment consisting of the following
- 15 approximate conditions:
- 16 72oF, and 50% humidity.
- The samples were weighed and then exposed to the
- 18 controlled environment. The samples were weighed before and
- 19 after exposure. The increase in weight is associated with
- 20 the increased moisture.
- The NaPBTC increased moisture through adsorbing
- 22 moisture from the surrounding environment.
- It is understood that based upon the known reactivity
- 24 of calcium hypochlorite, which is a Class 3 oxidizer, and

- 1 its known increased decomposition with increased moisture,
- 2 that the addition of an anti-scaling agent that increases
- 3 hygroscopicity will lead to a potentially dangerous
- 4 situation. Thus, the instant invention provides a
- 5 combination of ingredients which simultaneously reduces the
- 6 hygroscopic characteristics
- 7 of the final solid product, thereby improving the
- 8 manufacturing safety and final product stability, while
- 9 ensuring chemical feed reliability through improved deposit
- 10 control.

11 TABLE 1

12		Weight	Weight After	w/w % Change
13		Before	2hr	2hr
13	Dry Polymaleate	0.99grams	1.01grams	2.02%
14	Polymer	1 00	1 07	1.6 510
15 16	BAYHIBIT (PBTC)	1.09grams	1.27grams Weight	16.51% w/w % Change
			метдис	w/w % Change
			After 24hr	24hr
17	Dry Polymaleate		1.06grams	7.07%
18	Polymer			
19	BAYHIBIT (PBTC)		1.93grams	77.06%
20	Adsorption of la	ocalized mo	isture creates	the greatest
21	concern. While the	% weight in	crease to a 1.	5% blend would
22	appear to be minimal	, localized	moisture incr	ease will
23	correlate to the data	a exemplifi	ed in Table 1,	reducing shelf
24	life, chlorine activ	ity, and po	tential increa	sed risk of
25	combustion when fuel	sources are	e contacted as	outlined in
	McHale & Slavin, P.A.	15	519.023	8

- 1 Table 2, below.
- While not wishing to be bound to any particular
- 3 physical form, the combination of calcium hypochlorite and
- 4 polymer may be in the form of a homogeneous mixture, may
- 5 alternatively take the form of an outer layer or covering
- 6 which coats or encapsulates the cal-hypo for increased
- 7 stability and reduced reactivity, or a combination thereof.
- 8 Table 2 illustrates the reaction of 68% calcium
- 9 hypochlorite shock, sold under the brand name HTH SHOCK,
- 10 with and without a coating of a calcium polymaleate salt,
- 11 upon exposure to oil.
- Each sample used approximately 7.5 grams of solid
- 13 calcium hypochlorite. The treated sample was mixed with 4.7
- 14 grams of polymer salt (slurry weight), followed by
- 15 convective drying. Each sample was placed in a dish where
- 16 loz. of oil (brake fluid) was added as a fuel source.
- 17 The results clearly illustrate that coating the solid
- 18 calcium hypochlorite with a polymeric salt(s) of Polymaleic
- 19 acid or PESA as described in U.S. Patent 6,146,538
- 20 dramatically reduces the reactivity of calcium hypochlorite.
- 21 Even in the case of extended exposure to combustible
- 22 materials, the formulation described has the ability to
- 23 significantly reduce the rate of decomposition of the
- 24 calcium hypochlorite and prevent ignition when ignition

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1 would otherwise occur without treatment. It is therefore

2 also reasonable to extrapolate from the data that further

3 improvement would be achieved by encapsulation of the solid

4 calcium hypochlorite.

5 TABLE 2

6		Time to	Time to	
7	HTH Shock	Fuming 58	Combustion 62	Residue Charred, light
		seconds	seconds	weight, ash like
8	Treated HTH Shock	525	Not	residue No charred,
		seconds	Applicable	heavy, polymer
0	7 ddi+ion		formed mains	cal-hypo mix

Additional tests were performed using various polymeric salts with known anti-scaling abilities. These polymers all possess thermal stability sufficient to prevent their decomposition during exposure to exothermic reactions from the decomposition of exposed calcium hypochlorite as well as hot fuming fuels. The thermal stability ensures the polymer coating prevents exposure of coated calcium hypochlorite thereby preventing a self-sustaining decomposition or runaway reaction.

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1 Salt samples of polyacrylate and polymaleate were 2 produced for reactivity testing as illustrated in Table 3.

Table 3

4	Sample	Polymer Salt	Grams 68%	Grams Polymer		
<i>-</i>		Form	Cal-Hypo	(solids)		
5 6 7	1	Na/Ca Polyacrylate	45	Approx. 2		
8 9	2	Ca Polyacrylate	45	Approx. 2		
10 11	3	Na/Ca Polymaleate	45	Approx. 2		

These polymer salt samples were applied in either the form of a slurry or foam to 45 gram samples of >68% calcium hypochlorite sold under the brand name HTH. Foam (table 4 sample 3b) was used to enhance the distribution of the polymer. After drying, the treated samples were compared to blank samples for reactivity. 10ml of brake oil fluid was used as a fuel source, and the time from addition to fuming and ignition was measured, along with their weights.

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Table 4 illustrates the results of the testing.

Table 4 2

3	Sample	Cal-hypo wt.	Total wt.	Fuming (sec) Ig	gnition (sec)	Final Weight	Appearance	Polymer
4		(gm)	(gm)			(gm)		applied
4	Untreated							
5	1	7.72	9.99	spilled	n/a	n/a	n/a	n/a
6	2	7.84	10.02	12	14	10.13	Charred Ash	n/a
7	3	7.81	9.99	13	15	9.74	Charred Ash	n/a
8	4	7.86	10.03	13	15	9.37	Charred Ash	n/a
9								
10								
11								
12	Treated							
13		7.88	9.95	100	N/A	15.97	Cal-hypo residue	Slurry
14		7.77	10.03	105	N/A	16.8	Cal-hypo residue	Slurry
15		7.89	9.99	45	N/A	14.67	Cal-hypo residue	Slurry
16		7.88	9.92	195	N/A	18.83	Cal-hypo residue	Foam
177	шь			-h +	-	++ + h	+	na tha

17 The results of these test illustrate that treating the

>68% calcium hypochlorite with an alkali salt form of 18

hydrophilic polymer with thermal stability exceeding 300°F

significantly reduces the reactivity of the calcium

21 hypochlorite.

> The reaction is self-limiting in that only exposed (untreated) calcium hypochlorite is involved in the reaction and once consumed in the reaction, residual fuel remains in contact with the stabilized (treated) calcium hypochlorite.

These test results further illustrate that by limiting exposure of calcium hypochlorite to the fuel source, the rate of reaction is significantly reduced, and ignition is avoided all together.

Yet another result of these tests is that increased coverage of the calcium hypochlorite yields further McHale & Slavin, P.A.

- 1 reduction in reactivity of the treated calcium hypochlorite.
- 2 In this example, application of the polymer by use of an
- 3 expanded foam to increase its volume improved coverage and
- 4 distribution, correlating to improved performance at
- 5 reducing the samples reactivity.
- 6 As previously presented, improving the hygroscopic
- 7 characteristics of the calcium hypochlorite improves the
- 8 manufacturing safety and shelf-life of the solid product.
- 9 This is true regardless of whether the solid product is in a
- 10 granular, pellet, tablet or briquette form. By adding an
- 11 anti-scaling agent with improved hygroscopic
- 12 characteristics, the potential for compromising safety and
- 13 shelf-life is reduced.
- 14 By reducing the surface area of the calcium
- 15 hypochlorite by coating the solid material with a
- 16 hydrophilic polymeric salt with thermal stability exceeding
- 17 300oF, the reactivity of the calcium hypochlorite is
- 18 reduced, while allowing rapid dissolution of the calcium
- 19 hypochlorite when diluted with water, as in the case of a
- 20 calcium hypochlorite dispenser.
- 21 To further improve the hygroscopic characteristics of
- 22 the hydrophilic salt, divalent salts, or increased
- 23 percentages thereof, should be included in the final salt
- 24 form of the polymer.

- It is to be understood that while a certain form of the 1 invention is illustrated, it is not to be limited to the 2 specific form or arrangement of parts herein described and 3 It will be apparent to those skilled in the art that 4 shown. various changes may be made without departing from the scope 5 of the invention and the invention is not to be considered 6 limited to what is shown and described in the specification. One skilled in the art will readily appreciate that the 8 present invention is well adapted to carry out the objects 9
- 11 those inherent therein. The compounds, compositions, and any

and obtain the ends and advantages mentioned, as well as

- 12 biologically related compounds, methods, procedures and
- 13 techniques described herein are presently representative of
- 14 the preferred embodiments, are intended to be exemplary, and
- 15 are not intended as limitations on the scope. Changes
- 16 therein and other uses will occur to those skilled in the
- 17 art, which are encompassed within the spirit of the
- 18 invention and are defined by the scope of the appended
- 19 claims.
- 20 Although the invention has been described in connection
- 21 with specific preferred embodiments, it should be understood
- 22 that the invention as claimed should not be unduly limited
- 23 to such specific embodiments. Indeed, various modifications
- 24 of the described modes for carrying out the invention which

- 1 are obvious to those skilled in the art are intended to be
- 2 within the scope of the following claims.